KINETICS OF PRECIPITATE COARSENING IN A HIGH STRENGTH LOW ALLOY STEEL CONTAINING TI AND V

By MAHESH KUMAR AGARWAL

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KINETICS OF PRECIPITATE COARSENING IN A HIGH STRENGTH LOW ALLOY STEEL CONTAINING TI AND V

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MAHESH KUMAR AGARWAL

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CERTIFICATE

This is to certify that the present work, entitled Kinetics of Precipitate Coarsening in a High Strength Low Alloy Steel Containing Ti and V, by Mahesh Kumar Agarwal has been carried out under my supervision and it has not been submitted elsewhere for a degree

dm

April, 1993

Dr Shant P Gupta
Professor
Department of Metallurgical Engineering
Indian Institute of Technology
Kanpur

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April 1993 Yanpur Mahesh Kumar Agarwal

DEDICATED

TO MY

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ABSTRACT

The coarsening behaviour of carbonitride precipitate has been examined in a high strength low alloy steel (HSLA) containing 0 105 wt / Ti and 0 089 wt/ V as carbonitride forming elements in the temperature range 733-770°C steel was aged for periods ranging from 1 h to 343 h at 733°C. 752°C and 770°C temperatures Transmission Electron Microscopy was carried out on carbon extraction replica Particle size distributions (PSDs) were determined for various aging times at the three aging temperatures Although a = kt rate Law was observed during coarsening None of the presently available coarsening theories showed complete agreement with the experimental particle size A normal distribution rather than log-normal distribution was observed. The value of volume diffusivity, Dv is observed to be comparable with the corresponding published value

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LIST OF SYMBOLS

| Symbol | Meaning |
|-------------------|--|
| rxa B | Solute concentration in the matrix at the |
| °B | precipitation - matrix interface of a particle of |
| | r (Mole fraction) |
| ωχα B | Solute concentration in the matrix at the |
| B | precipitation - matrix interface of a large |
| | particle (Mole fraction) |
| Λ ^w (3 | Molar volume of the phase $(m^3/mole)$ |
| ນ | Interracial free energy of the 1th face |
| h | Time in hour |
| 7 | Temperature (°Y') |
| R | Gas constant (8 31434 J K^{-1} mole K^{-1}) |
| J ₁ | The flux to the particle from the matrix |
| J ₂ | The flux arising out of the growth of particles |
| r | Average particle radius at time t |
| 70 | Average particle at the onset of coarsening |
| v ^a | Coefficient of diffusion of the solute in the matrix |
| D | Rate constant |
| Q | Activation energy |
| dR | Growth rate |
| dt P | The ratio of the edge length to the average edge |
| 1* | length (a/a) |
| g (p) | Normalized distribution function |
| у (<i>г.</i> | Slope of line plotting inbetween $(\overline{a})^3$ vs t (m^3/sec) |
| | |

CHAPTER 1

INTRODUCTION

The mechanical properties of HSLA steel are strongly dependent upon the size and distribution of the carbide precipitate. Since the carbide particle can coarsen during the initial heat treatment and during subsequent service, It is important to be able to predict the kinetics of coarsening The ideas of precipitate coarsening was first formulated by Ostwald [1] as early as in 1900. The precipitate coarsening describes the change in average particle size with increasing time. It results from higher solubility of small particles, which may tend to dissolve in the matrix and cause the growth of larger particles The driving force is provided by the reduction in total interfacial energy of the system when the average particle size increases with time. This can be best illustrated with the help of the free energy-composition diagram, Figure 1.1 Let us consider a large spherical particle of radius r being surrounded by a number of smaller particles whose agerage radius is r. Figure 1.2 A common tangent construction between G_m^{α} and G_m^{β} phases of the free energy - composition diagram established equilibrium between the two phases Such construction indicates that large particles have lower solubility of the solute in the matrix in which they are contained. It is clear from the figure 1 1 that when large particles grow at the expense of smaller particles, there is a reduction in the interfacial energy per unit volume of the material. The energy thus released acts asdriving force for

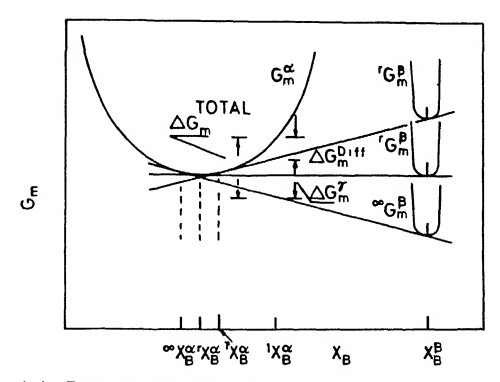


Fig 1 1 Free energy-composition diagram illustrating the driving force for coarsening

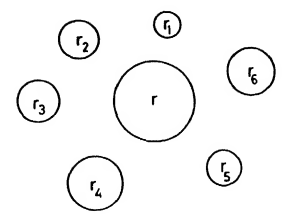


Fig 1 2 A polydisperse assembly of particles

the coarsening reaction

Because of the importance of the precipitate stability in commercial alloys the coarsening of precipitates has been studied experimentally by numerous investigators. Detailed theoretical analysis of the kinetics of coarsening was carried out by Lifschitz and Slyozov [2] and independently by Wagner [3] in what has become to be known as the LSW theory of coarsening. The principal finding of the LSW theory is that the average particle size increases with the cube root of the time of coarsening.

The solubility of a particle in the matrix as a function of its radius is best illustrated by the Gibbs - Thomson equation, written as

$$\ln \left[\Gamma_{X} \frac{\alpha / \beta}{B} / \infty_{X} \frac{\alpha / \beta}{B} \right] = \frac{\sqrt{\beta}}{RT} \frac{dA}{dV}$$
 (1 1)

where

 $r_{X_B}^{\alpha/\beta}$ = Solubility of a particle of radius r,

 ${}^{\infty}X_{B}^{\alpha/\beta}$ = Solubility of a particle of infinite radius,

 V_m^B = Molar Volume of the precipitate,

γ = Interfactal free energy.

 $\frac{dA}{dV}$ = Rate of change of surface area of the precipitate with volume and R & T have their usual meaning

For spherical particles of radius of n

$$A = 4\pi r^2$$
, $V = 4/3 \pi r^3$

So the term dA/dV in equation (1.1) equals 2/r which upon substitution yields

$$\ln \left[r_{X_B}^{\alpha/\beta} / {}^{\omega}_{X_B}^{\alpha/\beta} \right] = \frac{2V_m^{\beta} \gamma}{RTr}$$
 (1.2)

Assuming typical values of the terms on the right hand side of equation (1.2)

$$\gamma = 100 \text{ mJ/m}^2$$

R $\approx 8314\text{J/mole}^{\circ}\text{K}$

T = 1000°k
 $V_m^{\beta} = 10 \times 10^{-6}\text{m}^{3}/\text{mole}$

T = $10^{-7} \text{ to } 10^{-8}\text{m}$

Then

$$\frac{2\sqrt{\beta_{\gamma}}}{RTr} = \frac{2 \times 100 \times 10 \times 10^{-6} \times 10^{-3}}{8 \ 314 \times 1000 \times 10^{-7}} = 2 \ 4 \times 10^{-3}$$

Since the term on the right hand side has a small value, equation (1 2) is written as

$$\left(\Gamma \chi_{B}^{\alpha/\beta} - {}^{\omega}\chi_{B}^{\alpha/\beta} \right) = \frac{2V_{m}^{\beta}r^{-\omega}\chi_{B}^{\alpha/\beta}}{RTr}$$
(1.3)

If an isolated particle of the precipitate phase is rich in solute then the concentration distance profile in the early stages of its growth is as shown in Figure 1.3. However, nucleation of precipitate particles in the vicinity of an isolated particles and their continued growth make the diffusion field overlap. The particles stop growing thereafter. New region cannot nucleate solute-rich (3 phase because the neighbouring matrix is depleted of solute. The smaller particles dissolve during coarsening and the solute thus released moves down the concentration gradient (built gradually from smaller to larger particles) to attach themselves to larger particles. The processof coarsening therefore is one where some particles are continuously dissolving, some are continuously growing and some remain quasi-stationary at any given small time interval. During this interval some

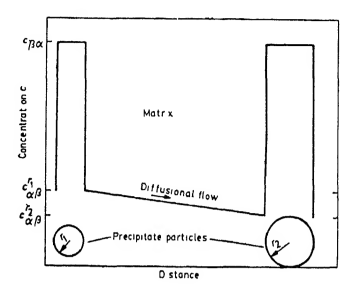


Fig 1 3 Concentration - distance profile on a particle which is growing

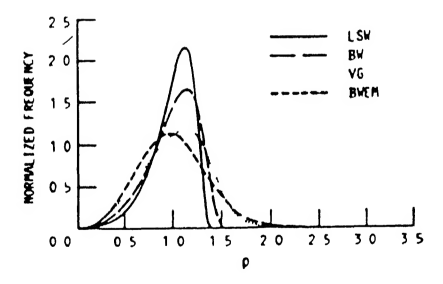


Fig 1 4 Theoretical particle distribution 51 Z e predicted by the LSW[5], BW[8], BWEM[9], The steady-state VG[10] theories distribution are presented for the BW and BWEM theories precipitate volume fraction of 60/ and for the VG theory at 50 Vol / [Ref 15]

particles must have dissolved completely and disappeared. The particles which remain quasi-stationary, i.e., which is neither growing nor dissolving at any given small time interval is termed as the critical size particle of radius r. It is understood that particles that remained quasi-stationary in a given interval may not be so in subsequent intervals. This is so because with the coarsening of particles, the average particle size increases and the critical particle size also increases with it. The net results being that the number of particles remaining after any time interval decreases with time and the average particle size increases with time.

There are a number of coarsening mechanism. These are

- 1 Volume diffusion controlled
- 2 Ledge mechanism
- 3 Grain boundary diffusion controlled
- 4 Diffusion through dislocations
- 5 Interface controlled

When the rate controlling mechanism is the diffusion of solute through the regular lattice sites, the coarsening mechanism is said to be volume diffusion controlled (VDC). In order to establish a functional relationship between the particle size and the time of coarsening, It is essential to make some simplified assumption. These are

The concentration gradient is considered to be time independent. This will allow us to use the Fick's first. Law to adequately describes the concentration gradient under steady state conditions.

- The solute diffusion coefficient, D is independent of concentration of the solute. This condition will again be fulfilled because dilute solutions are generally used in precipitation studies.
- The interfacial energy of the α/β interface $\nu_{\alpha\beta}$, is independent of the radius of curvature. This is quite critical below particle size of 100Å or so. Since the size of the particle at the termination of growth is of the order of 100Å, the interfacial energy will not be seriously affected during coarsening. Thus a constant value of $\nu_{\alpha\beta}$ will be assumed for the entire coarsening process.
- 4 For spherical particles, the effective diffusion distance is considered to be proportional to the radius of curvature of the particle. The value of the proportionality constant, α, which is reality is a complicated non-analytical function of several variables is assumed to remain constant during the entire coarsening process.

1 1 Volume Diffusion Controlled Coarsening

For spherical particles of radius r, the flux J_1 to the particle form the matrix is given by the Fick's first Law as

$$J_1 = -D 4\pi r^2 \frac{\Delta X_{\beta}^{\alpha}}{\Delta x} \tag{1.4}$$

Where D = rate of diffusion of solute through the lattice

 $4\pi r^2$ = surface area of the sphere

 $\Delta x =$ effective diffusion distance

From the Gibb ${\bf 3}$ Thomson ${\bf s}$ effect, equation (1.3), the solute concedintration difference $\Delta X_{\bf p}$ is given as

$$\Delta X_{B} = \begin{bmatrix} r \chi_{B}^{\alpha/\beta} - \overline{r} \chi_{B}^{\alpha/\beta} \end{bmatrix} = \omega_{X_{B}^{\alpha/\beta}} \frac{2\nu V_{m}^{\beta}}{RT} \begin{bmatrix} \frac{1}{r} - \frac{1}{r} \end{bmatrix}$$

and

Ax effective = ar

which upon substitution in equation (1 4) yields

$$J_{1} = -D 4\pi r^{2} \frac{\omega_{X_{B}}^{\alpha/\beta}}{\alpha RT_{r}^{2}} \left[1 - \frac{1}{r} \right]$$
 (1.5)

The flux J_1 can be equated to the flux J_2 arising out of the growth of spherical shaped particle from the matrix which gives

$$J_{2} = \left[X_{B}^{\beta} - X_{B}^{\alpha} \right] \frac{dV}{dt} = \left[X_{B}^{\beta} - \overline{X_{B}^{\alpha}} \right] 4\pi r^{2} \frac{dr}{dt} = J_{1}$$
 (16)

Where

V = Volume of the sphere and

. Upon equating the flux equation for J_1 and J_2 and rearranging terms, we get

$$\frac{dr}{dt} = -\frac{2DV_{m}^{\beta} \nu^{\alpha} \chi_{B}^{\alpha}}{\alpha RT \left(\chi_{B}^{\beta} - \chi_{B}^{\alpha}\right) r^{2}} \left[1 - \frac{r}{r}\right]$$
 (1.7)

On rearranging the terms of equation (1.7) and integrating on both sides we get

$$\int_{\overline{r}}^{\overline{r}} \frac{r^2 dr}{(1 - \frac{r}{r})} = -\int_{0}^{t} \frac{2D_{V} V_{M}^{\beta} v^{\omega} X_{B}^{\alpha}}{\alpha RT (X_{B}^{\beta} - X_{B}^{\alpha})} dt$$
(1.8)

Treating as constant and applying results of the LSW theory, equation (1 8) becomes

$$\vec{r} - \vec{r}_{D} = \frac{8}{9} D_{V} \frac{V_{D}^{\beta_{D}} \omega_{X}^{\beta/\alpha}}{\alpha RT (X_{B}^{\beta} - X_{B}^{\alpha})} t$$
 (1.9)

V

Where \overline{r} is the average particle radius after time t, \overline{r}_D is the average particle radius at the onset of coarsening. D_V is the diffusion coefficient of the solute in the matrix and other terms have already been defined

D Ramkrishna and S P Gupta (41 derived the equation for platelets shape particle by volume diffusion controlled coarsening. They consider square platelets of edge length a and thickness h. If the interfacial energies of the broad face and the edge with the α matrix are ν_F and ν_g respectively, application of the Gibbs Theomson equation yields

$$\ln \left(\frac{\chi_B^{\alpha}}{\omega_{\chi_B^{\alpha}}}\right) = \frac{\chi_B^{\beta}}{RT} \frac{4h\nu_e + 4a\nu_F}{2ah}$$
 (1 10)

Where X_B^α and X_B^α are the solubilities of particle of any size and a very large size respectively and V_m^β is the molar volume of the β phase

From the Gibbs-Wulff theorem, The following can be written

$$\frac{2\nu_{\rm F}}{h/2} = \frac{2\nu_{\rm e}}{a/2} = W \tag{1.11}$$

Where W is the Wulff constant, which on substitution in equation (1 11) gives the solubility difference between particles of edge length a and mean value \overline{a} as

$$\Delta X_{B}^{\alpha} = \frac{4 \nu_{a} V_{m}^{\beta \omega} X_{B}^{\alpha}}{RT} \left[\frac{1}{a} - \frac{1}{a} \right]$$
 (1 12)

The flux J_1 to the particle from the Fick's first Law is

$$J_1 = -D_v (2a^2 + 4ah) \frac{4 \nu_e v_m^{\beta} \omega_{XB}}{RT\alpha(a/2)a} \left\{ 1 - \frac{a}{a} \right\}$$
 (1.13)

Where $\alpha a/2$ is the diffusion distance, D_V is the volume diffusion coefficient and $2a^2+$ 4ah is the area through which diffusion takes place. The flux J_1 is equated to the flux J_2

arising out of the growth of particles as follows

$$J_{2} = \left(\chi_{B}^{\beta} - \chi_{B}^{\alpha} \right) \frac{d}{dt} \left\{ 4 \left(\frac{a}{2} \right)^{2} h \right\} = -D_{v} \left(2a^{2} + 4ah \right) \frac{4 \nu_{e} v_{M}^{\beta} \omega_{X}^{\alpha}}{R T \alpha a^{2} / 2} \left(1 - \frac{a}{a} \right)$$
(1 14)

Letting $h = \alpha/\beta$, differentiating and rearranging yields

$$\int_{a_{0}/2}^{a/2} \frac{(a/2)^{2} d (a/2)}{(1-\frac{a}{a})} = -\int_{0}^{t} \frac{2}{3} D_{v}(\beta + 2) \frac{\nu_{e} v_{B}^{\beta \omega} \chi_{B}^{\alpha}}{RT\alpha} dt$$
 (1.15)

The parameter β is related to the specific interfacial energies of the two interfaces and can be calculated. Treating α as a constant and applying results of the Lifshitz-Slyozov-Wagner theory gives

$$\left(\frac{\overline{a}}{2}\right)^{3} - \left(\frac{\overline{a}}{2}\right)^{3} = \frac{8}{27} (\beta + 2) D_{V} \frac{\nu_{e} V_{m}^{\beta \omega} \chi_{B}^{\alpha}}{\alpha RT (\chi_{B}^{\beta} - \chi_{B}^{\alpha})} t$$
 (1 16)

Except for the numerical factor 8/27 (β +2), equation (1.16) is identical in form with the kinetics equation of Wagner (3). Substitution for β = 1(h=a) enables the kinetics equation for the coarsening of cube-shaped precipitates to be obtained

$$\left(\frac{\bar{a}}{2}\right)^{3} - \left(\frac{\bar{a}}{2}\right)^{3} = \frac{8}{9} - \frac{D_{v}\nu_{e}v_{m}^{\beta\omega}\chi_{B}^{\alpha}}{\alpha RT \left(\chi_{B}^{\beta} - \chi_{B}^{\alpha}\right)} t$$
 (1 17)

$$(\bar{a})^3 - (\bar{a}_0)^3 = \frac{64}{9} \frac{D_v v_e v_m^{\beta \omega} \chi_B^{\alpha \alpha}}{\alpha RT (\chi_B^{\beta} - \chi_B^{\alpha})} t$$
 (1.18)

Equation (1 18) can be written in an approximate form

$$\bar{a} = (Kt)^{1/3}$$
 (1.19)

where

K(Rate constant) =
$$\frac{64}{9} \frac{D_{v} v_{em}^{\beta \omega} \chi_{B}^{\alpha}}{\alpha RT (\chi_{B}^{\beta} - \chi_{B}^{\alpha})}$$

1 2 Partical Size Distribution

Given sufficient time of coarsening of the precipitate after initial growth, particle which are larger than the average will grow and those smaller than the average size will gradually shrink and disappear. Thus a particle size distribution function, f(r,t) can be defined that corresponds to the number of particles of a given radius r at time t. The total number of particle n in the system at any instant is given by

$$n_p = \int_0^\infty f(r,t)dr$$

initially, n_p is constant and equals the number of particles at the end of the growth process of the precipitate. However, as some particles grow, some dissolve and others remain stationary during coarsening, the particle size distribution broadens. The distribution of particle sizes evolves towards a quasi-steady state distribution, f(r,t), which is independent of the initial distribution f(r,0). The distribution function is written as

$$f(r,t) = f(t) \rho^2 h(\rho) \qquad (1.20)$$

Where f(t) is function of time only

$$\rho = \frac{r}{r}$$

and $h(\rho)$ as solved by Wagner for a Gaussian distribution is

$$h(\rho) \left(\frac{3}{3+\rho}\right)^{7/3} \left(\frac{3/2}{\frac{3}{2}-\rho}\right)^{11/3} \exp^{-\left(\frac{\rho}{3/2-\rho}\right)}$$
 (1.21)

From which

$$h(\rho) = 0 \quad \text{for } \rho = \rho_{\text{m}} = 3/2$$

 ρ_m being one cut-off radius ratio also known as the maximum relative radius of the particle. Taking $f(r,t)\alpha$ $\rho^2h(\rho)$, it is clear that,

for
$$\rho = 0$$
, $f(r,t) = 0$
and for $\rho = 3/2$ $f(r,t) = 0$

This implies a sharp cut-off in th distribution, such that no precipitate particles exists with radius more than 1.5 times the mean radius of the particles. The distribution has a maximum value at $\rho=1.135$

Following Wagner's approach [3], the integration can be carried out for

$$\rho = \frac{r}{r^*} = \frac{r}{r} = \frac{3}{2} \text{ yielding}$$

$$\int_{\frac{3}{2^{r}}}^{\frac{3}{2^{r}}} \frac{r^{2} dr}{1 - \frac{3}{2}} = \frac{2D \nu_{\alpha\beta} V_{m}^{\beta\omega} \chi_{B}^{\alpha}}{\alpha RT \left(\chi_{B}^{\beta} - \chi_{B}^{\alpha}\right)} t$$
(1 22)

Or

$$(\overline{r})^3 - (\overline{r}_0)^3 = \frac{8}{9} \frac{D_v \nu_{\alpha\beta} V_m^{\beta\omega} \chi_B^{\alpha}}{\alpha RT (\chi_B^{\beta} - \chi_B^{\alpha})} t$$
 (1 23)

Although the cube rate law has been exhibited by a number of alloys. The experimental particle size distribution (PSD s) do not concide with the prediction of the LSW [5] theory. This discrepancy is mainly attributed to the fact that the LSW theory is strickly applicable when the precipitate volume fraction is small and approaches zero.

approximation is not valid for many material systems. As a result, considerable effort has gone into modifying the basic theory. For example, Lifshiz-Slyozou Encounter Modified (LSEM) theory. It has been shown by Davies et al. [7] which considers encounter between precipitate particles, the Brailsford-Wynblatt (BW) [8], the Brailisford-Wynblatt Encounter Modified (BWEM). [9], and the Voorhees-Glicksman (VG) [10] theories.

The theoretical PSD s predicted by the LSW [5], the BW [8] BWEM [9], and the VG [10], theories are shown in Figure 1.4. The latter three theories have attempted to incorporate the influence of finite precipitate volume fraction. Figure 1.4 shows the steady state distribution which are predicted by the BW and BWEM theories for alloys with 60 vol / precipitate and by the VG theory for alloys with 50 Vol/ precipitate. In the present study, the experimental PSD s were determined as a function of time and temperature.

1 3 Effect of Volume Fraction of the Prcipitate

The theory of particle coarsening by Lifshitz and Sigozov and Wagner (LSW), is applicable only when the volume fraction of the precipitate is very small, under the condition, the mean distance between particles is large compared to the dimension of the particle. Since the kinetics of coarsening is controlled by the diffusion of solute from a particle which is dissolving to a particle which is growing, it is expected that the mean distance between particles which is controlled in turn by the volume fraction, V_f of the precipitate will influence the rate of growth As V_f increases, the mean separation between particles decreases and the distance that solute atoms have to diffuse on an average

to become a part of the growing precipitate become shorter. A detailed analysis of the problem has been treated by Ardell [5] by considering a distribution of particle sizes surrounding an ith particle of radius r_1 , as shown in Figure 1.5. If the mean free path between the ith particle and its neighbour is \overline{X} then, for a poly dispersed assembly of particles, the distance r at which the solute concentration in the matrix approaches the solute concentration of the particle – matrix interface of average size particle is given by

$$r = r + \frac{\overline{\chi}}{2}$$
 (1 24)

In order to introduce the effect of the volume fraction on the rate of coarsening, a relation between r and $V_{\rm f}$ must be obtained. This relation is difficult to obtain for a polydisperse assembly. However, when the particles are considered to be all of the same radius r^* , the center to center distance, d, between a particle and its nearest neighbour is

$$d = r^* \left\{ 2 + \frac{e^{8V_f}}{3^3 \gamma_{V_f}} (V_f) \right\}$$
 (1.25)

where the gamma function,

$$\Gamma(V_f) = \int_{BV_f}^{\infty} Y^{-2/3} e^{-y} dy$$

Now, from figure 1 5, $\overline{X} = d - 2r^*$

$$= \frac{e^{BV_f}}{3^3 V_f} (V_f) r^*$$
 (1 26)

and
$$r = r + \frac{r^*}{2} - \frac{8V_f}{3^3 v_f} + T(V_f)$$
 (1.27)

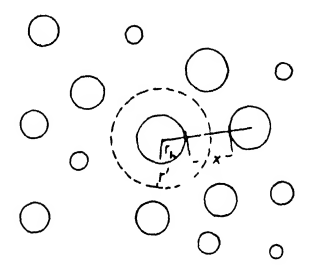


Fig 1 5 A polydisperse assembly of particles showing the mean free path

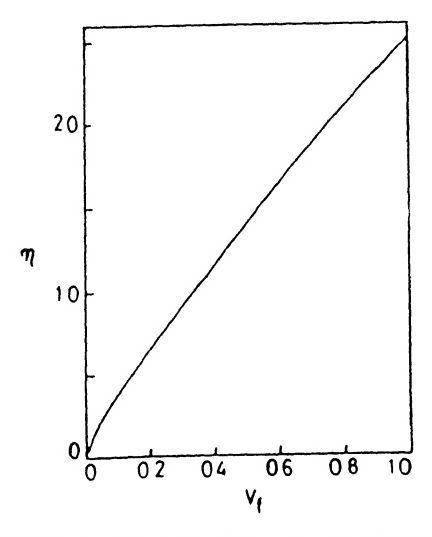


Fig. 1.6 The effect of the volume fraction on the parameter η

or
$$\frac{\Gamma}{\Gamma} = 1 + \frac{1}{\eta P}$$
 (1 28)

where $\eta = \frac{2 \ 3^3 / v_f}{e^{BV} f(v_c)}$ and $P = \frac{\Gamma}{r}$

The effect of volume fraction on the parameter η is shown in Figure 1.6

Rate equation which is as follows

$$\frac{dr^{3}}{dt} = \frac{6D \nu_{\alpha\beta} V_{m}^{\beta} v_{B}^{\alpha}}{RT (X_{B}^{\beta} - X_{B}^{\alpha})} (1 + \eta P) (\rho-1)$$

or,

$$r^{*3} - r^{*3} = \frac{6D \nu_{\alpha\beta} V_{m}^{\beta} \omega_{XB}^{\alpha}}{RT (X_{B}^{\beta} - X_{B}^{\alpha}) \psi} t$$
 (1.29)

Where r_{Ω}^{*} is the critical radius of the particle in the initial distribution

r * is critical size which is neither growing nor dissolving

$$\psi = \frac{4D \nu_{\alpha\beta} V_{m}^{\beta} \omega_{XB}^{\alpha}}{RT (X_{B}^{\beta} - X_{B}^{\alpha})} \frac{dt}{dr^{*3}}$$

 $\frac{\overline{\rho}}{\rho} = \frac{\overline{r}}{r}$ which upon substitution in equation (1.14)

$$(\overline{r})^{3} - (\overline{r}_{\alpha})^{3} = \frac{6D \nu_{\alpha\beta} v_{m}^{\beta} \omega_{XB}^{\alpha}}{RT (X_{B}^{\beta} - X_{B}^{\alpha}) \psi} (\overline{\rho})^{3} t \qquad (1.30)$$

The value of $\overline{\rho}$ has been calculated by Ardell [5] and it is found to have weak dependence on the volume fraction of the precipitate. In the range that most studied of coarsening are carried out, $\overline{\rho}$ increases from approximately 0.94 to 1.0 as the volume fraction decrease from 0.1 to zero.

1 4 Estimation of Q amd D

Equation (1 19) is of the form $y = K^{1/3}X$ where $K^{1/3}$ indicate the slope of line. From Plot of \overline{a} vs. $t^{1/3}$ atdifferent temperatures and Dv can be calculated. The Arrhenius equation is

$$D_{v} = D_{q} \exp^{-Q/RT}$$
 (1.31)

where D_v is the diffusion rate, D_o is the rate constant

Q is the activation energy and

R&T have their usual meaning

Upon substitution of the value of D in equation (1 19) we get

$$K = \frac{64}{9} \frac{D_{\alpha} v \sqrt{\beta} \propto \chi_{B}^{\alpha}}{\alpha RT \left(\chi_{B}^{\beta} - \chi_{B}^{\alpha}\right)} \exp^{-Q/RT}$$
 (1 32)

This can be written as

$$KT = \frac{64}{9} \frac{D_0 \nu V_m^{\beta} \omega \chi_B^{\alpha}}{\alpha R (X_B^{\beta} - X_B^{\alpha})} * xp^{-Q/RT}$$
 (1 33)

or KT = A exp^{-0/RT}

or
$$\ln VT = \ln A - \frac{Q}{RT}$$

where

$$A = \frac{64}{9} \frac{D_{\alpha} \nu \ V_{m}^{\beta} \ \omega_{X_{B}^{\alpha}}}{\omega_{R} \ (X_{B}^{\beta} - X_{B}^{\alpha})}$$

We can determine the diffusion parameters D_0 and Q from the intercept and slope of the straight line respectively from the Arrhenius plot. On compairing the theoretical and experimental values of D_0 and Q one can predict the mechanism that is operating for the growth of precipitates

CHAPTER 2

EXPERIMENTAL PROCEDURE

21 Material

The present investigation was carried out to study the kinetics of precipitate coarsening in a HSLA Steel containing. To and V. The HSLA steel containing 0.1/ Vanadium and 0.08/ Titanium was obtained in the hot forged condition from Climax. Molybdenum Corporation U.S.A. Table 2.1 shows the composition of this steel in the as - received condition. It contains small amount of nitrogen which makes Vanadium nitride and Titanium nitrite precipitation possible during isothermal aging. The elements Si and Al are in very minor quantity and thus will have negligible influence. The sole function of these two elements will be in contributing to solid solution strengthening. The addition of Mn retards the rate of growth of ferrite from Austenite.

The steel had been hot forged to obtain rounds 18 mm in diameter and then cold rolled with a number of intermediate annealing treatments to obtain section 3 mm thick. Small square specimen size 6x6mm² were solution treated for 8 minutes Isothermal transformation was carried out at temperatures 733°C, 752°C and 770°C in sealed quartz tube under vacuum for 1 h to 15 days

The experimental procedure and the techniques used for the study of kinetics of precipitate coarsening are as described in the following sections

Table 2 1
Composition of HSLA Steel (in \text{ \text{\text{\text{Composition}}}}

| Element | С | N | Sı | Al | Mn | ٧ | Tı |
|---------|-----|-------|------|-------|-----|-------|-------|
| Amount | 0 1 | 0 016 | 0 09 | 0 048 | 1 5 | 0 089 | 0 105 |

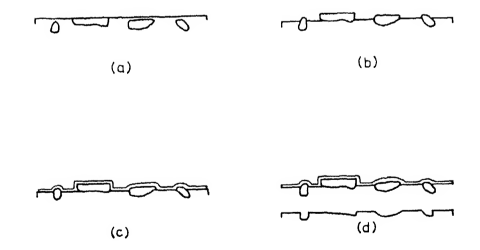


Fig 2.1 The extraction replica technique (a) Polished surface (b) Etched to expose second phase, (c) Carbon coated, and (d) re-etched and replica floated off [Ref 11]

2 2 Replica Preparation

In order to study the kinetics of precipitate coarsening of TiVCN the particle size and shapes were derived from carbon extraction replica technique This technique is illustrated schematically in Figure 2 1 The two phase structure to be replicated was first etched sufficiently in a nital solution containing 3/ nitric acid to expose the second phase layer of carbon coating was done in the usual way Shadowing of carbon is to be avoided in order to ensure a strong continuous film and therefore, rotation of the specimen or diffuse deposition Subsequently shadowing was used to give information about the three dimensional shape of the extracted particles specimen was next dipped in a nital solution containing 12/ nitric acid so that the matrix was attacked while the second phase, which was to be extracted remained unchanged. The replica, containing particles of the second phase was collected on a copper grid most critical part of this procedure was the second etching to free the replica and particles from the base metal It was necessary to score the carbon coating into squares before etching This floating carbon film containing particles was collected on copper grid (dia = 3mm) for further study

2 3 Transmission Electron Microscopic Studies

Carbon extraction replicas collected on copper grid were examined using the TEM (JEOL-2000FX). In order to confirm the shape and size of the precipitate sufficient number of photographs were taken and in order to confirm precipitate phase is indeed Vanadium Titanium carbonitride, selected area diffraction patterns were obtained from isolated large particle.

2 4 Particle Size Measurement

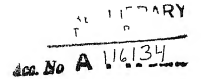
Edge length of the particle was measured manually with the help of a vernier caliper, having least count 0 002cm. From each samples measurements were made on approximately 500 particles

2 5 Actual Magnification Calibration

The electron photomicrographs taken on 35 mm camera don t have magnification marker on them. In order to carryout the calibration the same features were recorded on both the 35 mm camera and plate camera at a number of magnifications. The plate camera had the magnification marker on it. The same precipitate particle size is measured in both the photograph, and the actual magnification of 35 mm camera determined.

26 Particle Size Distribution

The particle size distribution data were obtained by feeding the raw data to a computer programmed to give mean particle size, size distribution and particle size ratio, a/a for each specimen



CHAPTER 3

RESULTS AND DISCUSSION

3 1 Morphology

A photomicrograph of the precipitate during coarsening is shown in Figure 3 1, which is a transmission electron micrograph of extraction replica derived from a specimen aged for 64 hours at 770°C clearly shows the square cross section of It precipitate In order to confirm it, the precipitate phase indeed Vanadium-Titanium carbonitride (VTiCN), selected diffraction pattern was obtained from isolated large particle as shown in Figure 3.2, as an illustration It indexes centered cubic (FCC) lattice. The cube geometry was maintained for the entire duration of coarsening as confirmed from the TEM micrograph of Figure 3.3 derived from a specimen aged for 216 hours at 752°C During later stages of coarsening, there was slight tendency for the edges of the cube to either get rounded or convert into small facets. With these modifications, the cube shaped particles appeared as slightly rounded as shown in Figure 3 4, derived from a specimen aged for 343 hours at 733°C The square geometry was not eliminated, however

3 2 Coarsening Behaviour of Precipitate:

The coarsening behaviour of the VTiCN precipitate was examined as a function of temperature from 733°C to 770°C for the 0 081 wty V, 0 0105 wt/ Ti, 0 1Wt/ C and 0 016Wt/ N containing HSLA Steel The TEM photomicrographs of Figure 3.5 depict the change in the microstructure which occurs during aging at 770°C

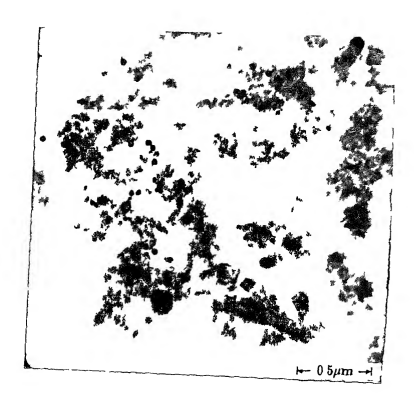


Fig 3 1 TEM Micrograph of extraction replica, specimen aged for 64 hours at 770°C

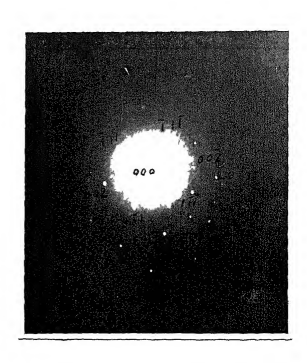


Fig 3 2 Electron diffraction pattern derived from a isolated precipitate particle

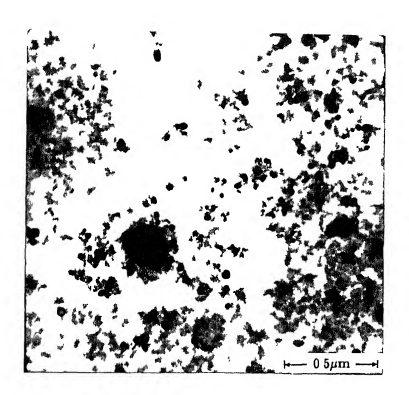


Fig 3 3 TEM Micrograph showing vanadium Titanium Carbonitride as cube geometry, specimen aged 216 hours at 752°C

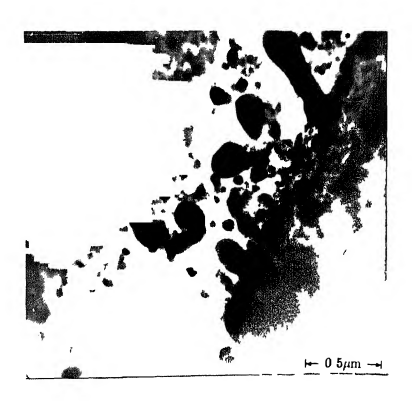


Fig 3 4 In TEM Micrograph Cube Shaped particles appeared as slightly rounded

for times of up to 216 hours Figure 35(a) shows the micro structure after aging for B minutes The precipitate particles in this condition were cubical 10 shape and had average particle size, $\bar{a} = 78.22$ angstrom After aging for 1h, 8h. 27h, 64h and 216h the precipitate particles increase in size, but some were coalescing as well Figure 3 5(b) derived from specimen aged for 216 hours at 770°C shows the increased average particle size

Similar results were obtained during coarsening at 752°C and 733°C. The precipitate particle were cubical in shape as shown in Figures 3 6(a) and 3 6(b) derived from specimens aged for 8 hours and 64 hours respectively at 752°C. Photomicrographs presented in Figures 3 7(a) and 3 7(b) show the precipitate particle derived from a specimen aged for 8 minutes and 64 hours respectively at 733°C.

3 3 Particle Size Distributions:

The initial particle size distribution after 8 minutes—aging at 770° C is shown in Figure 3.8(a). The aging time was sufficient for complete—transformation—of—austenite—to—ferrite—at—this temperature—The austenite—to—ferrite—transformation—initiates after an incubation period of few seconds and it is expected—that the Vanadium—Titanium—Carbonitride particle—that formed first must have coarsened to some extent—in—8 minutes—This cannot—be avoided, however, in this alloy system in which the second—phase forms by interface controlled mechanism—[12]—The coordinates—of Figure 3.8(a) are $g(\rho)$ and ρ —Where $\rho(a/\overline{a})$ is the—ratio—of—the edge length—to—the average edge length—and $g(\rho)$ is the—normalized distribution function, i.e., the fraction of—the—particles—that

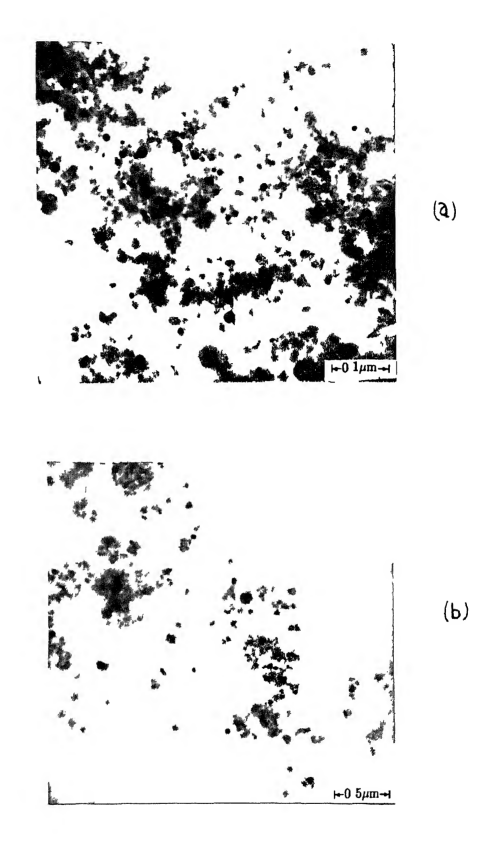


Fig 3.5 Transmission Electron Micrograph specimen aged at 770°C (a) aged for 8 minutes (b) aged for 216 hours

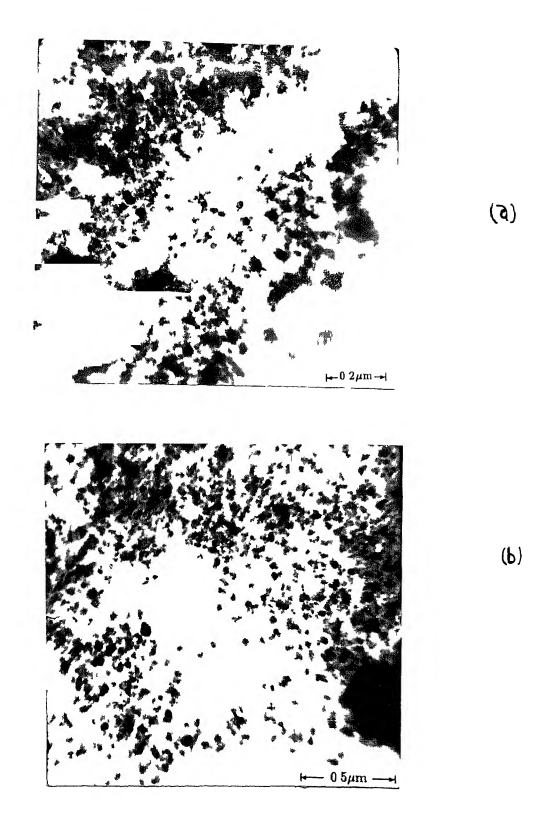


Fig 3.6 Transmission electron micrograph specimen aged at 752°C (a) aged for 8 hours (b) aged for specimen 64 hours

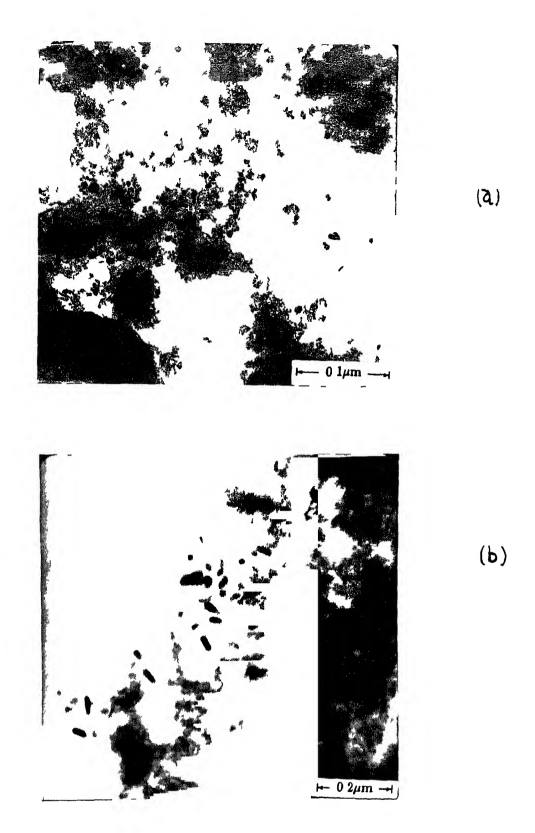
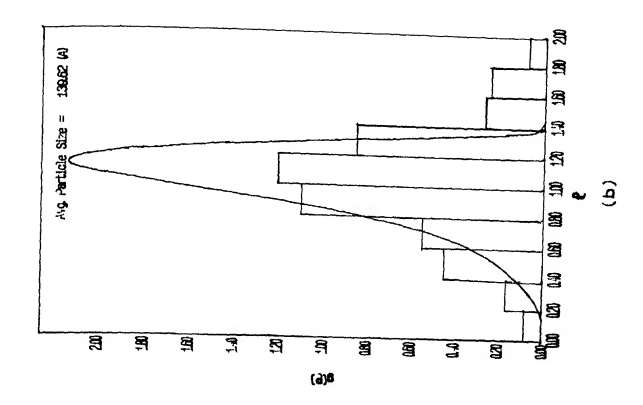


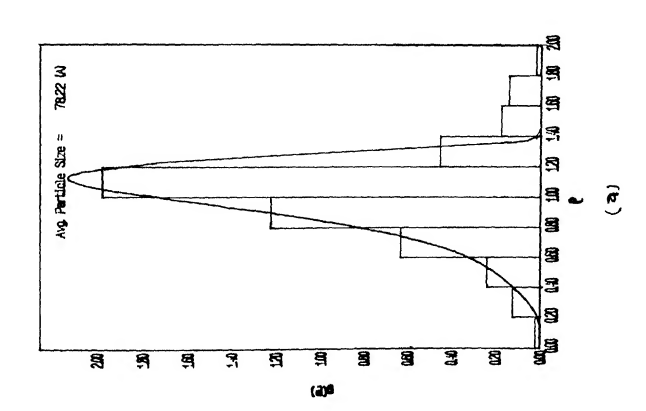
Fig 3 7 Transmission Electron Micrograph specimen aged at 733°C (a) aged for 8 minutes (b) aged for 64 hours

lie in an interval multiplied by the reciprocal of the interval

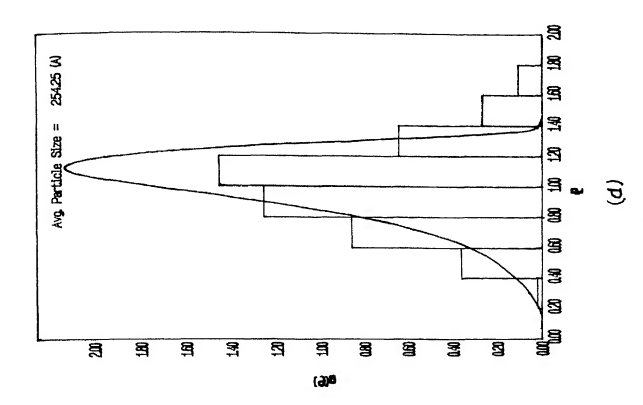
The particle size distributions after aging for 1h, 8h, 27h, 64h, 216h at 770°C are displayed in Figure 3 B(b), 3 B(c), 3 B(d), and 3 B(e) respectively. Values of ρ at which $g(\rho)$ is maximum lie in the range of 1 2 to 2 0 and the cut off radii ratio, ρ_{max} , lie in the range of 1 8to 2 0. PSDs after coarsening for 8 minutes 1h, 8h, 64h, 137h, 216h and 343h at 752°C are shown in Figure 3 9(a) - 3 9(g) respectively. PSDs for 8 minutes, 1 h, 64 h, 216h and 343 h at 733°C are shown in Figure 3 10(a)-3 10(e) respectively. Values of ρ at which $g(\rho)$ is maximum lie in the range of 1 0 to 1 8 and their cut off radii ratio ρ_{max} , lie in the range of 1 B-2 0. Based on the particle size distribution displayed in Figure 3 B to 3 10 at three different temperature, the following observations can be made

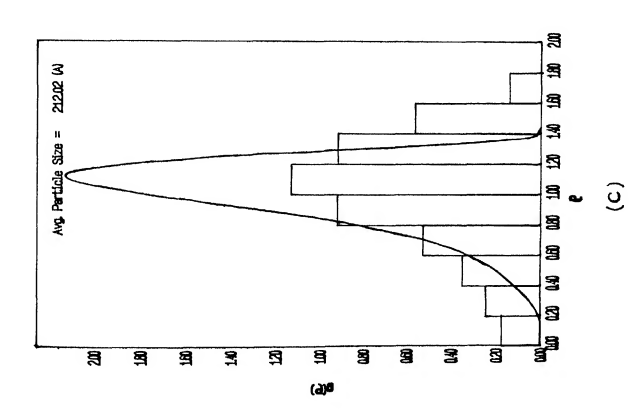
- (a) The particle size distributions for most aging condition were symmetrical about the mean value, indicating a normal distribution rather than a log normal distribution predicted by the Lifshiftz, Slyozov and Wagner [2,3] theory. The coarsening of VC in a Vanadium bearing HSLA Steel is reported (4) to have similar PSDs
- (b) The cut off diameter ratio, a/a is reported to have value upto 1 9 instead of the cut off radius ratio ρ of 1 5 as predicted by LSW [2,3] theory for the volume diffusion controlled coarsening. A cut off radius ratio exceeding. 1 5 has also been observed in a number of investigation, namely Ni-Co-Al [7], Al-Cu [13] and Fe-C-V [4].
- (c) The peak value of $g(\rho)$ is observed to fall in the range



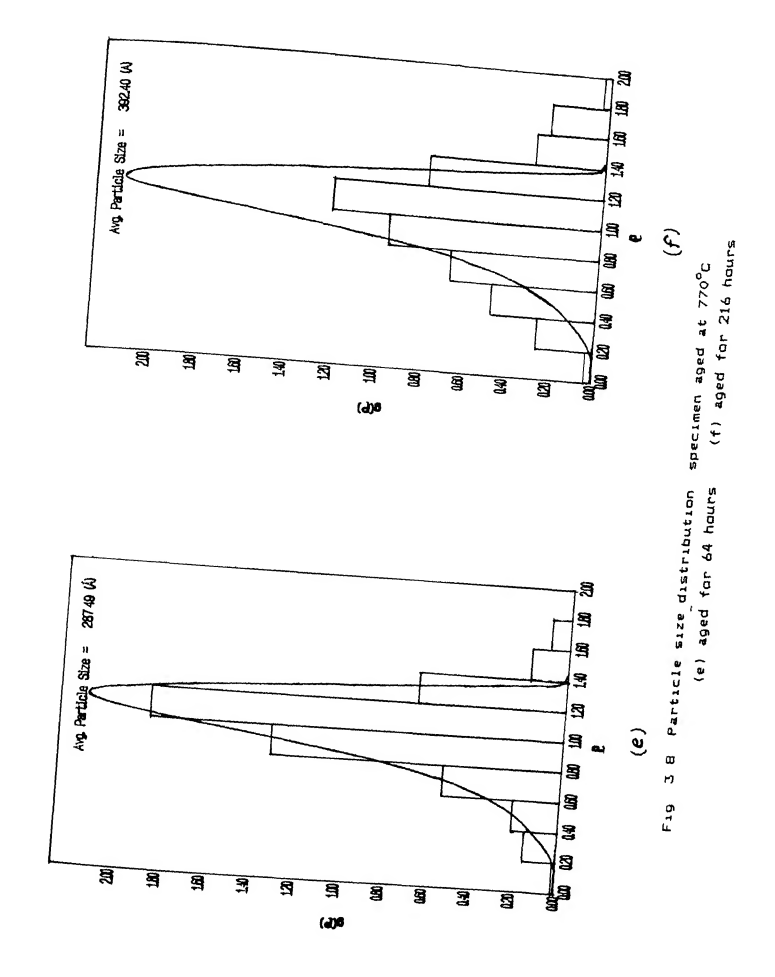


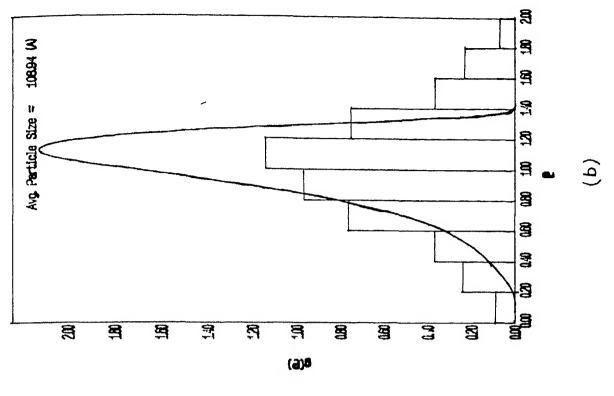
Particle size distribution, specimen aged at 770°C (a) aged for B minutes (b) aged for 1 hour œ ۲) F19

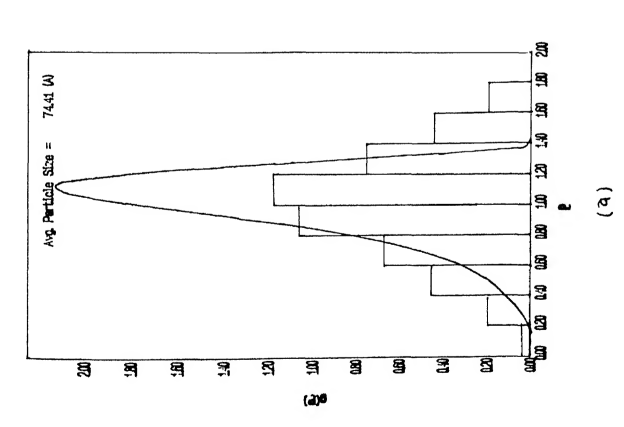




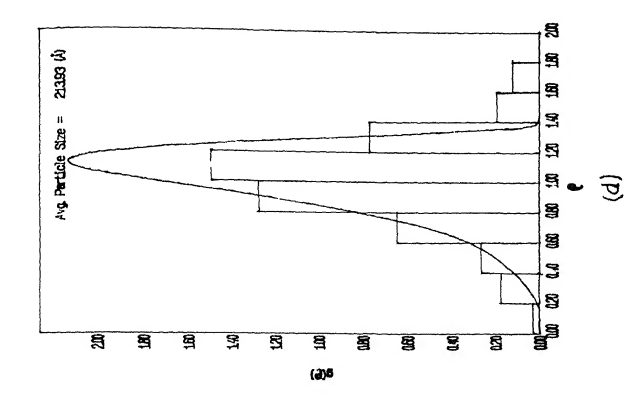
Particle size distribution, specimen aged at 770°C (d) aged for 27 hours (c) aged for 8 hours 63 CH

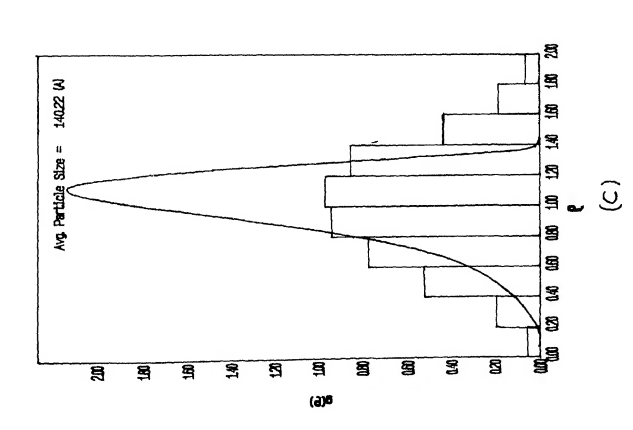




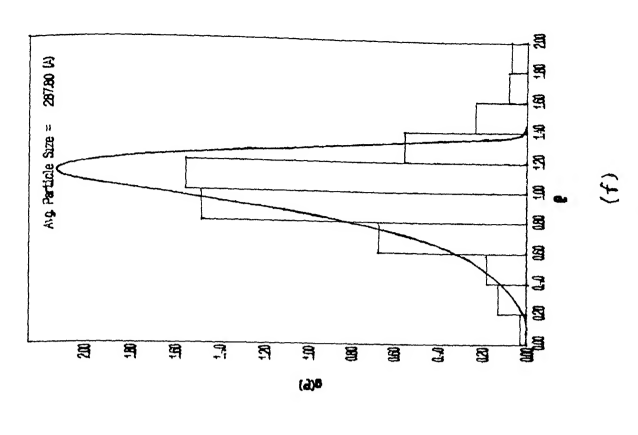


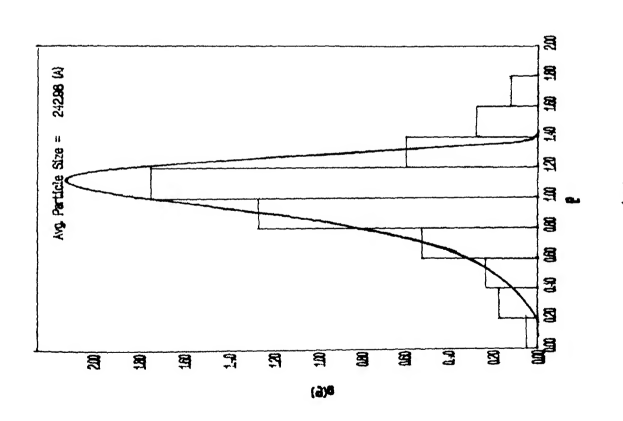
Particle size distribution specimen aged at 752°C (a) Aged for 8 minutes (b) Aged for 1 hour F19 39



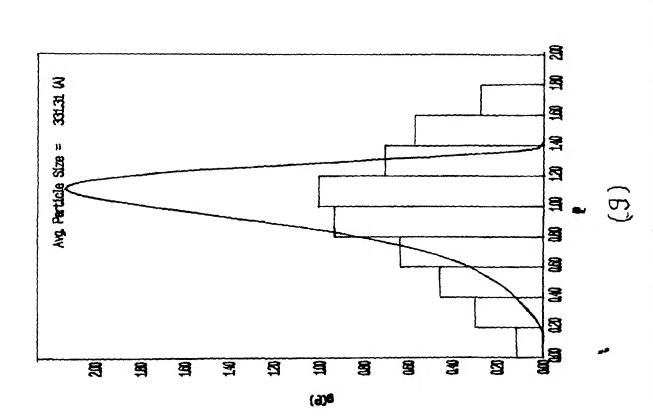


752°C (d) Aged for 64 hours J 9 Particle size distribution specimen aged at (c) Aged for 8 hours F19



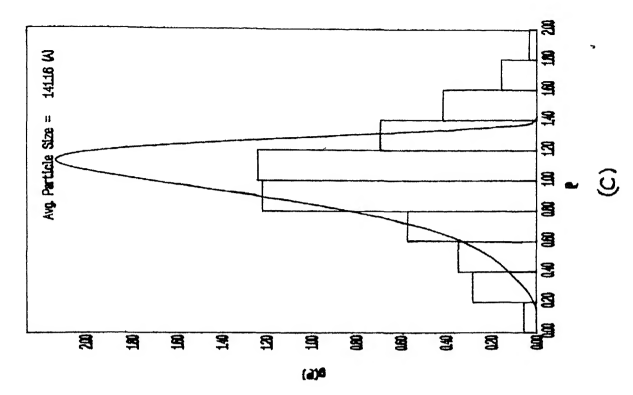


 $_{
m 3}$ 9 Particle size distribution specimen aged at $_{
m 752}^{
m o}$ C (e) Aged for 137 hours (f) Aged for 216 hours F19



Particle size distribution specimen aged at 7 3 10 3 10 Particle size distribution specimen aged Fig (g) Aged for 343 hours at 752°C 51,

(a) aged for 8 minutes



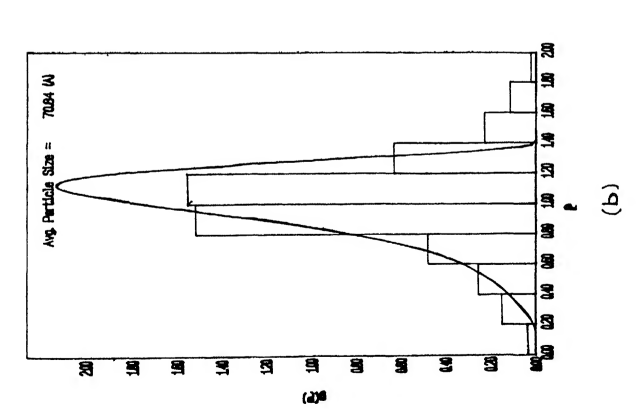
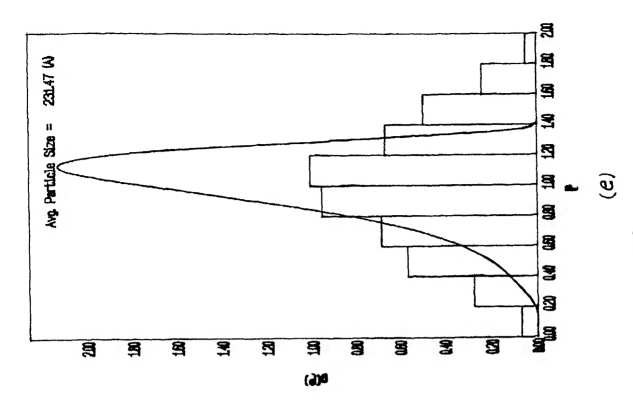
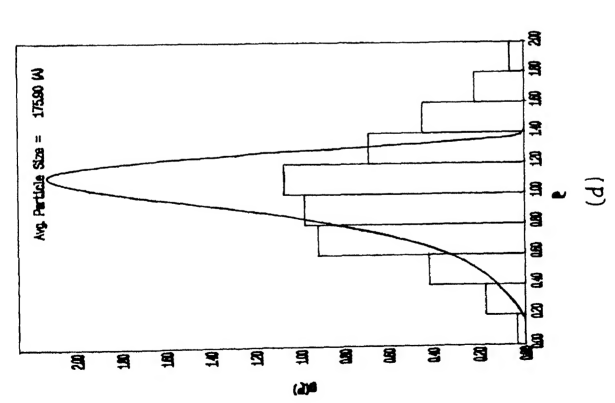


Fig 3 10 Particle size distribution specimen aged at 733°C (c) Aged for 64 hours (b) Aged for 1 hour





3 10 Particle size distribution specimen aged at 733°C (e) Aged for 343 hours Aged for 216 hours (p) F19

- 1 O-1 4 which is lower than the theoretically predicted value of 2 12 for LSW theory. Exception are observed in few cases where the peak value lies between 1 6 and 1 8
- (d) Instead of being narrow, a number of PSDs were broad near the value as displayed in Figure 3.9(c) and 3.10(d) after coarsening for 8 h at 752°C and 216 h at 733°C respectively

The above observation for the coarsening of VTiCN are consistent with those reported for vanadium nitride precipitates [14] and vanadium carbide [4]. Recently Mackay et al. [15] have shown that the PSCs during volume diffusion controlled coarsening of the phase in Nickel base alloys is a normal rather than a log normal distribution.

In an analysis carried out to study the effect of the volume fraction on the particle size distribution, Ardell [5] has shown that the peak value of g(p) decreases with increasing volume fraction ϕ of the precipitate and the PSD broaden. The broadening is symmetrical about value [Fig 3 11) At very large volume fraction of the precipitate, the PSD for volume diffusion controlled coarsening reaches that for interface controlled coarsening. The PSD, however remains asymmetrical as in the The Lifshiz-Slyozov encounter modified [LSEM] [2,3] theory theory of Davies et al [7] predicts a normal distribution symmetrical about the mean value and is particularly applicable for alloy system with a small volume fraction of the precipitate The basic kinetics equation for the growth remains (Fig 3 12) unchanged

If the LSEM theory has any validity then one must observe encounter between particles, i.e., coalescence of two or more

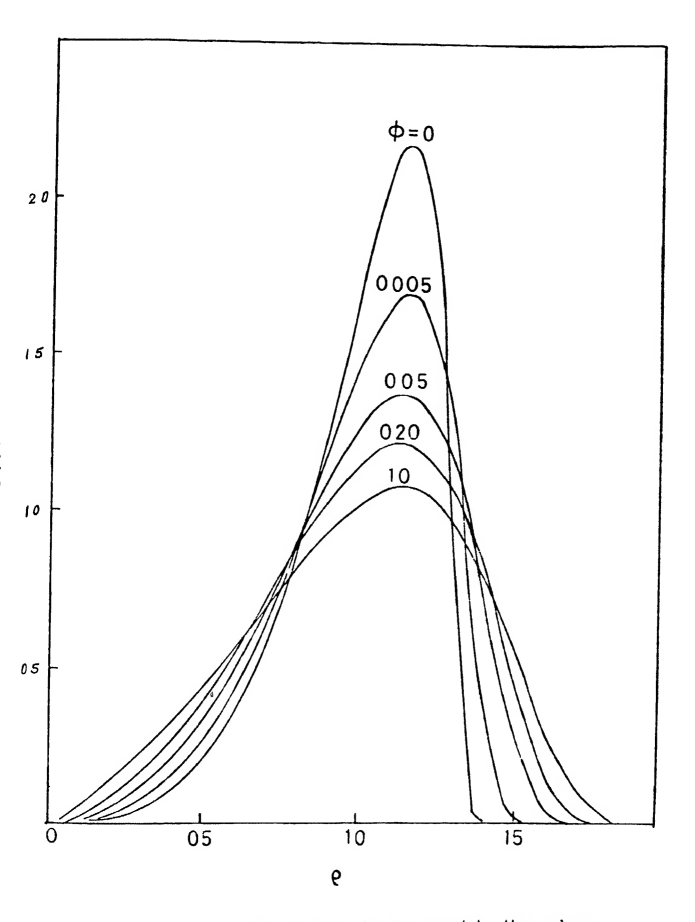


Fig 3 11 Theoretical PSD modified to accommodate the volume fraction of the precipitate [Ref 5]

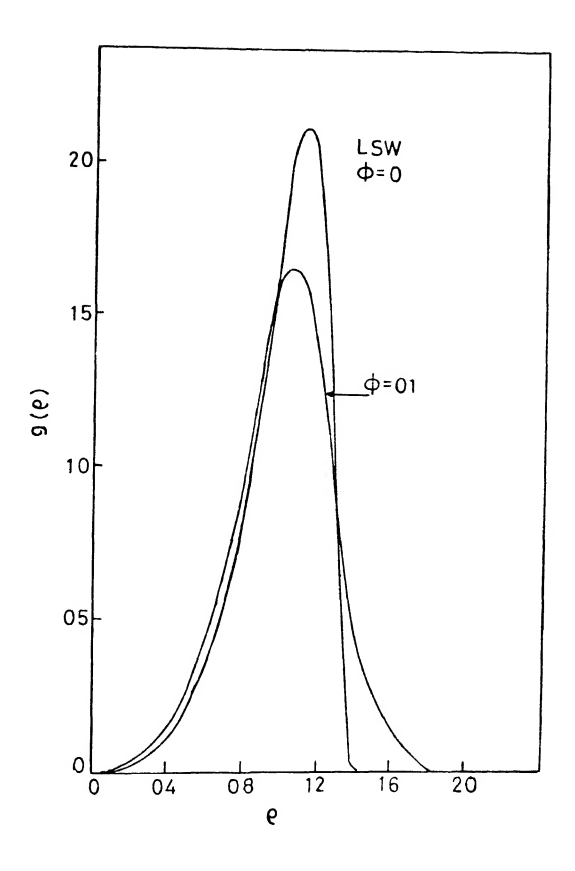


Fig 3 12 Theoretical PSD modified to incorporate the encounters between particles [Ref 7]

neighbouring particles which coarsened by receiving solute from the matrix. During the course of this investigation we observed coalescence between particles in a few specimen. The encounter were between two or more particles in the TEM micrograph of Figure 3.7(b). In some cases the particles were simply lying one above the other. Davies et al. [7] have shown L shaped and a number of rectangular shaped particles of Ni_3Al after 44 hours of aging at $750^{\circ}C$ in a Ni-Co-Al alloy of the LSEM theory of Davies et al. [7] is applicable. It will,

- (1) Modify the PSD so as to be symmetrical about the mean value
- (11) Increase the cut-off radius ratio to value close to 1 9 or more
- (111) Broaden the PSD near the mean value, and
- (IV) Lower the peak value of g(p)

Although the particle encounter theory of Davies et al [7]have some merit consistent with the observation both micro structural as well as of particle size distribution, contributes significantly only when the volume fraction vanadium Titanium Carbonitride in HSLA Steels containing comparable amount of V, Ti, C and N is reported to be VTiCN does not nucleate homogeneously but nucleates on coherent v/a interface during ferrite transformation by the ledge mechanism [17,18] Generally, a large step (ledge height) followed by several much smaller steps which leads to several closely spaced bands of precipitate followed by a large space to the next band of precipitate [18] Thus, a large volume fraction of the precipitate is confined within a sheet In pure Fe-V-C alloys, an intersect spacing of the order of 200 to 1000 Å

reported [17] Because of the large volume fraction of the precipitate, encounters between particles within a sheet and between particles within a row are expected. From the shape of the PSDs and micro structural evidence, the encounter modified theory of Davies et al. [7] deserves some merit

3 3 Kinetics of Coarsening

The kinetics equation describing time dependence of the particle size is derived for spherical or plate shaped particles. The Vanadium Titanium Carbonitride (VTiCN) precipitate is cube shaped assumed to have interfacial free energies of the coherent face ν . The equation derived by Ramakrishna et al. [4] is for plate shaped particle. In their equation (1.16) β = 1 (length/thickness) is replaced for cube shape particle, the basic equations for volume and interface diffusion controlled coarsening are identical in form to those of the LSW theory. For volume diffusion controlled coarsening these are

$$\left\{\frac{\overline{a}}{2}\right\}^{3} = \left\{\frac{\overline{a}_{0}}{2}\right\}^{3} = \frac{8}{9} \frac{D_{v} \nu_{g} V_{m}^{\beta} w_{g}^{\alpha}}{\alpha RT \left(X_{B}^{\beta} X_{B}^{\alpha}\right)} + (1.20)$$

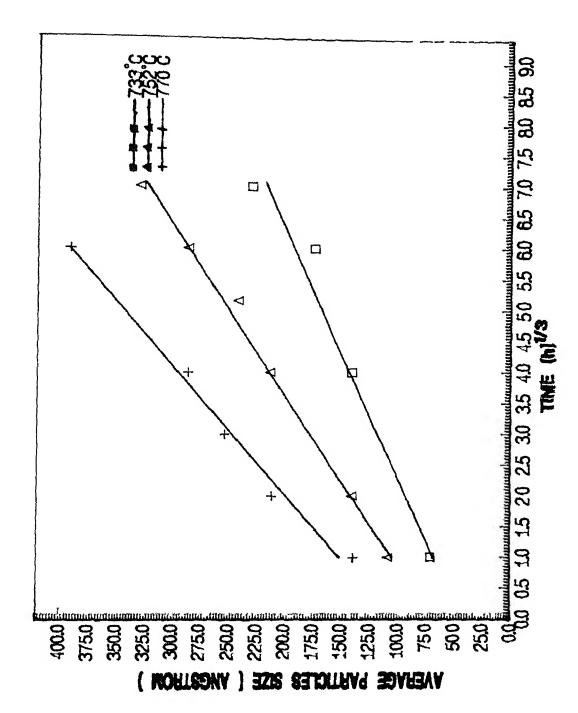
Taking the approximate form of the above equation, the average value of the particle size, \overline{a} (\overline{A}) is plotted against cube root of the time (t) $^{1/3}$ at the three temperatures used in this study. Figure 3-13 clearly shows that the average particle size data fits the t $^{1/3}$ proportionality. The experimental data for volume diffusion controlled growth are shown in Table 3-1

The coarsening kinetics therefore appears to be volume diffusion controlled as also reported for Fe-Mo-C (20), Fe-Ti-Si and Fe-V-C (4) alloys. The data of Figure 3 13 were fitted

Table 3 1

Average Particle Size with respect to Time and Temperature

| Temperature | Aging Time | Average Particle Size |
|-------------|------------|-----------------------|
| (°C) | (h) | A |
| 770 | 1 | 137 62 |
| 770 | 8 | 212 02 |
| 770 | 27 | 254 25 |
| 770 | 64 | 287 49 |
| 770 | 216 | 392 40 |
| 752 | i | 108 94 |
| 752 | 8 | 140 22 |
| 752 | 64 | 213 93 |
| 752 | 137 | 242 98 |
| 752 | 216 | 287 80 |
| 752 | 343 | 331 31 |
| 733 | 1 | 70 84 |
| 733 | 64 | 141 16 |
| 733 | 216 | 175 90 |
| 733 | 343 | 231 47 |
| | | |



3 13 a versus ${\rm t}^{1/3}$ plot for the coarsening of vanadium Titanium Carbonitride (VTICN) in HSLA Steel F19

to linear least square constant (\overline{a}_0) and slope $K^{1/3}$, thus determined are shown in Table 3.2

In an attempt to get some idea of the atomic species controlling the rate of coarsening, the equation (1.18) was used and calculation of the diffusion coefficient $D_{_{V}}$ made at the three temperatures. The constant Y in this equation is dependent upon the interfacial energy of the edge with the α matrix, the parameter α and β , mole fraction of solute in the α phase in equilibrium with a very large size precipitate, ${}^{\infty}X_{\beta}^{\alpha}$, the molar volume of the vanadium-titanium carbonitride (VTiCN) and the difference $(X_{\rm R}^{\beta} - X_{\rm R}^{\alpha})$

From the crystal structure (FCC) and the published lattice parameters which falls in the range 4 138 to 4 182Å for VC_{O 75} to VC_{O 961} and in the range 4 3127-4 328Å for $T_1C_{O 61} - T_1C_{1 O}$ Taking the approximate value of the lattice parameter to be 4 25Å for T_1VCN It gives the value of V_M^β to be 11 558 × $10^{-6} \, \mathrm{m}^3/\mathrm{mole}$ For $^\infty X_B^\alpha$, we used B= carbon and determined the mole fraction of carbon in ferrite at various temperatures from the equilibrium Fe-C phase diagram. This is approximate only. The exact value can be obtained from the $\alpha/(\alpha+VT_1C)$ boundary in a steel containing 0 185VT1 and 1 50 Mn. Since X_B^α is very small compared to X_B^β the difference $(X_B^\beta-X_B^\alpha)$ is taken to be \approx 0 45 for VT1CN. The parameter α is treated as a constant with a value equal to unity. α is not a constant but is dependent upon the particle size as shown by Shiflet et al. [221]

The interfacial free energy as the coherent broad face is expected to be low and comparable to the coherent twin boundary energy of 30 mJ/m 2 [23]. On this basis alone the interfacial

Table 3 2 Value of the Constant (\bar{a}_0) and Slope $(N^{1/3})$

| Temperature | a _o | r ^{1/3} | r |
|-------------|----------------|-----------------------|-----------------------------------|
| °C | (Å) | (A/h ^{1/3}) | (m ³ s ⁻¹) |
| 770 | 78 22 | 46 26 | 27 5 x 10 ⁻³⁰ |
| 752 | 74 41 | 36 67 | 13 78 × 10 ⁻³⁰ |
| 733 | 54 69 | 23 48 | 3 6 × 10 ⁻³⁰ |
| | | | |

ı

Table 3 3 $\text{Values of K, } \overset{\omega}{\times} \overset{\alpha}{B}, \text{ } D_{_{\mathbf{Y}}} \text{ experimental and Calculated }$

| | yers, 10-11-11-11-11-11-11-11-11-11-11-11-11-1 | | |
|--|--|-------------------------|--------------------------|
| Temperature | 770°C | 752°C | 733°C |
| K, m ⁵ | 27 5×10 ⁻³⁰ | 13 78×10 ⁻³⁰ | 3 6×10 ⁻³⁰ |
| ωχα B | 1 486×10 ⁻³ | 1 72 x 10 ⁻³ | 2 135×10 ⁻³ |
| D _v , Experimental m ² s-1 | 1 952×10 ⁻¹⁸ | 0 830×10 ⁻¹⁸ | O 171×10 ⁻¹⁸ |
| D _v , m ² s ⁻¹ Ramakrishana et al [4] | 0 16x10 ⁻¹⁸ | *0 09×10 ⁻¹⁸ | **0 05x10 ⁻¹⁸ |
| D _v , m ² s ⁻¹ Bowen et al [25] | 282×10 ⁻¹⁸ | *168x10 ⁻¹⁸ | ** _{92×10} -18 |
| D _v , m ² s ⁻¹ Bowen et al [2 6] | 2 7×10 ⁻¹⁸ | *1 4×10 ⁻¹⁸ | **0 8×10 ⁻¹⁸ |
| D _v , m ² s ⁻¹ Moll et al [27] | 111×10 ⁻¹⁸ | 67×10 ⁻¹⁸ | 12×10 ⁻¹⁸ |
| 0 | | | |

^{*} at 750°C

^{**} at 730°C

energy of the edge can be computed to be about 110 mJ/m². The average energy of the edge can be taken as approximately 250 mJ/m² which is lower than the grain boundary energy $(700\text{mJ/m}^2 \text{ of ferrite}$ and interfacial energy of 700mJ/m^2 for the Fe/Fe₃C interface [24] at 710°C

The value of $D_{_{V}}$ was calculated from approximate values of the parameters in equation (1.18) as shown in Table 3.3. The diffusivity values of this investigation are comparable to the $D_{_{V}}$ values obtained by Ramakrishna et al. (V in α -Iron) [4], which is very much close to value obtained by Bowen et al. (V in α -Iron) [25] and the experimental $D_{_{V}}$ values differ from those obtained by Bowen et al. (V in α -Iron) [26] and Moll et al. (Ti in α -Iron) [27] by an order of magnitude two. The data compare very favourably with the experimentally observed $D_{_{V}}$ values. From these $D_{_{V}}$ values at different temperature, activation energy (Q) and $D_{_{O}}$ can be calculated. However, due to insufficient points the calculations of $Q_{_{V}}$ and $D_{_{O}}$ were not made

CHAPTER 4

CONCLUSIONS

The rate of growth of Titanium Vanadium Carbonitride in HSLA Steel exhibited cube rate law behaviour and valume diffusion controlled mechanism predicted by the LSW theory of particle coarsening. The LSW theory as well as several modified theories were ineffective in describing the experimentally determined particle size distribution experimentally observed PSDs follow a normal distribution rather than a log-normal distribution There is some evidence of the modification of the PSDs by encounters between particles However, the exact contribution they make in modifying the PSDs is not known The experimental particle size distribution differ with respect to the peak value, cut off radius ratio and shape of predicted PSDs from theoretical considerations estimated value of the parameter of the kinetics equation, the experimental diffusivity, $D_{_{\mathrm{U}}}$, values are comparable with the calculated $\mathbf{D}_{_{\mathbf{U}}}$ values as they lie within an order of magnitude of each other

REFERENCES

- 1 W Ostwald, Z Phys Chem (Leipzig), 34 (1900), p 495
- 2 I M Lifshiz and V V Slyozov, J Phys Chem Solids, 19 (1961), p 35
- 3 C Wagner, Z Electrochem, 65 (1961), p 61
- 4 D Ramakrishna and S P Gupta, 92 (1987), p 179 Mat SC Elriff
- 5 A J Ardell, Acta Metallurgica, 20 (1972), p 61
- 6 W Greenwood, Acta Metallurgica, 4 (1956), p 243
- 7 CKL Davies, P Nash and R N Stevens, Acta Metallurgica, 28 (1980), p 179
- 8 A D Brailsford and P Wynblatt Acta Metallurgica, 27 (1979), p 489
- 9 CS Jayanth and P Nash, J Mater Sci , 24 (1989), p 3041
- 10 PW Voorhees and ME Glicsmon, Acta Metallurgica, 32 (1984), p 2013
- 11 P J Goodhew, Specimen preparation in Material Science,
 Newyork, (1973)
- 12 J M Gray and R B G Yeo, Trans ASM, 61 (1968), p 255
- 13 P Merle and F Fouquet, Acta Metallurgica, 29 (1981), p
- 14 O E Atasoy, Metall Trans A, 14 (1983), p 379
- 15 R A Mackay and M V Nathal, Acta Metallurgica, vol 38, 6 (1990), p 993
- 16 A D Batte and R W K Honeycombe, Met Sci, J, 7 (1973), p
 160
- 17 KR Kinsman and HI Aaronson, Met Trans , 4 (1973), p
 959

- 18 R W K Honeycombe, Metall Trans A , (1976), p 915
- 19 NK Balliger and RWY Honeycomb, Met Trans A, 11A (1980), p 421
- 20 D M Davies and B Ralph, J, J Iron Steel Inst , London, 210 (1972), p 262
- 21 E Bower and J A Whitemann, The Mechanism of Phase
 Transformation in Crystalline Solids, Institute of Metals,
 London, (1968), p. 119
- 22 C J Shiflet and H I Aaronson, Acta Metallurgica, 27 (1979), p 377
- 23 M Mclean and H Mykura, Acta Metallurgica, 12 (1964), p
 326
- 24 J J Kramer, 6 M Pound and R F Mehl, Acta Metallurgica, 6 (1958), p 763
- 25 A W Bowen and G M Leak, Metall Trans , 1 (1970), p 2767
- 26 A W Bowen and G M Leak, Metall Trans 6 (1970), p 1965
- 27 S H Moll and R E Ogilvie, Trans Metall Soc, AIME, 215 (1959), p 673

APPENDIX

```
COMPUTER PROGRAM FOR PARTICLE SIZE DISTRIBUTION
#irclud
#includ (str n h)
#include (rath h)
          hrrmle_name[18]
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                    0 j ( rumber_of_particles[i] 1 j++)
              (1
          fsc rf(itp /f &size[i][j])
                                                                                             ************
                                     CALCULATE THE REAL SIZE OF THE PARTICLES
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1 *
          real_fx e[i][j] = s e[i][j]*1 0e+07/magnxtude_of_magnxf[x]
          dumny_ 1 e[n] eal_size[i][j]
           ++
          3
          fclose(ifr)
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                    * * * *
11
              * *
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m > size (re l_s e[0][0] > real_size[0][1]) ? real_size[0][0] real_size[0][1]

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INITIALIZATION

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         fclose(1tp)
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